## **Amendments to the Claims**

This listing of claims will replace all prior versions, and listings, of claims in the application:

## **Listing of Claims:**

- 1. (Original) A method for the identification and investigation of a receptor in target tissue for which a selected vector has affinity, said method comprising:
  - i) creating retroviral particles containing a library of mRNA from the target tissue;
  - ii) transfecting a non-adherent cell line which does not bind with the selected vector by infecting the cells with said retroviral particles;
  - adding to the transfected cell line a suspension of encapsulated gas microbubbles to which the selected vector is coupled and allowing the microbubbles and cells coupled thereto to float to the surface of the suspension;
  - iv) isolating the microbubble-bound cells at the surface;and either
  - v-a) Iysing the isolated cells, amplifying the receptor-encoding cDNA therefrom and sequencing said cDNA; and optionally
  - v-b) comparing the thus-obtained sequence data with gene bank sequence data;

or

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- vi-a) culturing the isolated cells; and
- vi-b) investigating affinities of vectors to the isolated cells.
- 2. (previously amended) The method according to claim 1 wherein said vector is selected from peptides, proteins, antibodies, nucleotides, hormones, growth factors, cytokines, carbohydrates, lipids, therapeutic agents and drugs acting through receptor-mediated cell entry.
- 3. (previously amended) The method according to claim 1 wherein the encapsulated microbubbles of step iii) are selected from microbubbles of gas stabilised by a coalescence-resistant surface membrane, a filmogenic protein, a polymer material, a lipid, a non-polymeric and non-polymerisable wall-forming material and a surfactant.

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- 4. (previously amended) The method according to claim 3 wherein said surfactant is selected from one or more phospholipids and one or more lipopeptides.
- 5. (previously amended) The method according to claim 1 wherein said gas is a biocompatible gas or gas mixture selected from perfluorinated gases, preferably from sulphur hexafluoride, perfluoropropane, perfluorobutanes, perfluoropentanes and perfluorobexanes.
- 6. (previously amended) The method according to claim 1 wherein said gas is perfluorobutane and said surfactant is phosphatidylserine.
- 7. (previously amended) The method according to claim 1 wherein the microbubbles are removed before or after culturing, said removal is effected by bursting with a technique selected from ultrasonication, pH change or transient application of overpressure or underpressure.
- 8. (Original) Microbubble-bound transfected cells producible by method steps i) to iv) of claim 1.
- (Original) Microbubble-bound transfected cells according to claim 8 wherein the
  microbubbles are of similar size to the transfected cells, preferably the microbubbles have diameters of 1
  to 10 um, more preferably 3 to 5 um.
- 10. (cancel) Use of microbubble-bound cells according to claim 8 for the investigation of diseases involving said receptors.